11 Publication number:

**0 264 231** A1

(2)

## **EUROPEAN PATENT APPLICATION**

21 Application number: 87308942.9

(1) Int. Cl.4: C07D 205/08, A61K 31/395

2 Date of filing: 09.10.87

3 Priority: 17.10.86 JP 246638/86

49 Date of publication of application: 20.04.88 Bulletin 88/16

Designated Contracting States:
 AT BE CH DE FR GB IT LI LU NL SE

 Applicant: TAISHO PHARMACEUTICAL CO. LTD
 24-1 Takata 3-chome Toshima-ku Tokyo 171(JP)

② Inventor: Kawashima, Yutaka 1731-1, Akoda Tatebayasi-shi(JP)

Inventor: Satoh, Masakazu

Ekimae Puraza 6-205 15-1, Akamidai-2-chome

Konosu-shi(JP)
Inventor: Hatada, Yuichi

8-17, Minamimagome-4-chome Ota-ku

Tokyo(JP)

Inventor: Hazato, Fumiko

Kopo Sanraizu 203 41-7, Haraichi

Ageo-Shi(JP)

Inventor: Nakashima, Yoshimoto

18-16, Gobancho Ageo-shi(JP) Inventor: Sota, Kaoru 1158-11, Shimotomi Tokorozawa-shi(JP)

Representative: Ellis, Edward Lovell et al MEWBURN ELLIS & CO. 2/3 Cursitor Street London EC4A 1BQ(GB)

- Azetidinone derivatives.
- 2-Azetidinone derivatives represented by the following formula

P 0 264 231 A1

wherein X is a hydrog n atom, a halogen atom, a lower alkyl group, a lower alkoxy group, a hydroxyl group or a cyano group, t is 1 or 2, R¹ is a lower alkyl group, a cycloalkyl group, a 1-naphthylmethyl group, an

optionally substitut d phenethyl group, an optionally substitut d phenyl group, an optionally substituted benzyl group or a bis(alkoxycarbonyl)ethyl group, and R² is a lower alkyl group, a lower alkoxy group, an amino group, an adamantyl group, a lower alkoxycarbonylmethyl group or an optionally substituted phenyl group, are useful as blood platelet aggregation inhibiting agents.

### **AZETIDINONE DERIVATIVES**

#### BACKGROUND OF THE INVENTION

#### 1. FIELD OF THE INVENTION

The present invention relates to 2-azetidinone derivatives having blood platelet aggregation inhibiting activity.

#### 2. DESCRIPTION OF THE PRIOR ART

Although some compounds having azetidinone skeleton which show antibacterial activity have been known in the past, any azetidinone derivative showing blood platelet aggregation inhibiting activity has not been yet reported.

### SUMMARY OF THE INVENTION

15

As a result of earnest researches to blood platelet aggregation inhibiting activity of the compounds having an azetidinone skeleton, the present inventors have found novel 2-azetidinone derivatives having blood platelet aggregation inhibiting activity, and the present invention has been completed.

An object of the present invention is to provide 2-azetidinone derivatives represented by the general formula

$$\mathbb{R}^{2} \longrightarrow \mathbb{R}^{2}$$

$$0 \qquad \mathbb{R}^{2}$$

$$0 \qquad \mathbb{R}^{2}$$

$$0 \qquad \mathbb{R}^{2}$$

wherein X is a hydrogen atom, a halogen atom, a lower alkyl group, a lower alkoxy group, a hydroxyl group or a cyano group, L is 1 or 2, R¹ is a lower alkyl group, a cycloalkyl group, a 1-naphthylmethyl group, a 1-phenethyl group, 1-carboxy-2-phenethyl group, a group of the formula

(wherein Y and Y' are the same or different and each is a hydrogen atom, a halogen atom, a lower alkyl group, a lower alkoxy group, a dimethylamino group, a carboxyl group, a dichloroacetyl group or a trifluoromethyl group, or Y and Y' together form a methylenedioxy group, and m is 0 or 1) or a group of the formula

(wherein R<sup>3</sup> is a lower alkyl group), and R<sup>2</sup> is a lower alkyl group, a lower alkoxy group, an amino group, an adamantyl group, a lower alkoxycarbonylmethyl group, a group of the formula

(wherein Z is a hydrogen atom, a halogen atom, a lower alkyl group, a lower alkoxy group, a phenyl group or a nitro group, and n is 1 or 2).

Other object of the present invention is to provide blood platelet aggregation inhibiting agents containing the compound of formula I.

### DETAILED DESCRIPTION OF THE INVENTION

5

15

30

35

45

In the present invention, the term "lower alkyl group" refers to straight or branched chain alkyl group having 1 to 4 carbon atoms such as, for example, a methyl group, an ethyl group, a proppyl group, an isopropyl group, a butyl group, an isobutyl group, a tert-butyl group and the like. The term "cycloalkyl group" refers to a cyclopentyl group and a cyclohexyl group. The term "lower alkoxy group" refers to those having 1 to 3 carbon atoms such as, for example, a methoxy group, an ethoxy group, a propoxy group and the like. The term "halogen atom" refers to a fluorine atom, a chlorine atom, a bromine atom and an iodine atom. The term "lower alkoxycarbonylmethyl group" refers to those such as, for example, a methoxycarbonylmethyl group, an ethoxycarbonylmethyl group and the like.

Preferred compounds of formula I are those wherein X is a hydrogen atom, R<sup>1</sup> is a benzyl group or a chlorobenzyl group, and R<sup>2</sup> is a nitrophenyl group.

The compounds of the present invention can be easily prepared, for example, by a reaction (i.e., Wittig Reaction) of a compound represented by the general formula

wherein R1, X and I are as defined above, with a Wittig reagent represented by the general formula

wherein R2 is as defined above.

Reaction solvents used in this reaction are those used in the ordinary Wittig Reaction such as, for example, benzene, ethyl ether, tetrahydrofuran, toluene, chloroform, methylene chloride, dimethoxyethane and the like. The reaction temperature is from -30°C to the temperature of the boiling point of the solvent used, preferably from 0°C to 30°C. The reaction time depends on the starting material, the Wittig reagent or the reaction temperature, but usually it is from 2 to 48 hours, and the reaction may be stopped after the disappearance of the starting material observed by using thin layer silica gel column chromatography.

Configuration of the oxyalkylidene substituent of especially useful compounds of the present invention is E-form, and the configuration due to the asymmetric carbon atom at the 4-configuration is di-form.

Som of the compounds of formula II are known, and some are new and can be pr pared by the methods described in the literature [e.g., Tetrahedron Letters, Vol. 25 (No. 42), page 4733 (1984)].

It is recognized that the compounds of the pres nt invention have xcellent blood platel t aggregation inhibiting activity and v ry poor bleeding tendency as side-effect, and therefore, they are useful as blood platelet aggregation inhibiting agents. For the purpose, these compounds can be administered orally or parenterally in a conventional dosage form such as tablets, powders, granules, capsules, solutions, emulsions, suspensions, injectional solutions and the like, each of which can be prepared by conventional pharmac utical practices.

The dosage used as blood plat let aggregation inhibiting agents to human depends on the age, weight or response of patient, administration route or time of administration, but usually it may be from 10 to 3000 mg per day.

The LD<sub>∞</sub> of the compound of formula I in mouse is more than 5000 mg/kg.

Next, the following experiments illustrate concretely excellent blood platelet aggregation inhibiting activity and prolongation effect of bleeding time of the compound of the present invention.

#### 5 Experiment 1 [invitro test in rabbit]

Citrated blood (one volume of 3.2% sodium citrate; 9 volumes of blood) was collected from carotid artery of male, New Zealand strain house rabbit, centrifuged at 150 g for 15 minutes to give platelet rich plasma (PRP) as a supernatant, and the remaining blood was centrifuged at 1500 g for 10 minutes to give platelet poor plasma (PPP) as a supernatant. The platelet count of PRP was adjusted to  $50 - 60 \times 10^4 \mu l$  by dilution of PPP. Blood platelet aggregation was carried out according to the method of Born [Born, G.V.R., Nature, 194, 927 (1962)]. Namely, 25  $\mu$  of the test drug, (all the test drugs were dissolved in dimethyl sulfoxide and adjusted to the desired concentration with physiological saline solution), was added to 250  $\mu$  of PRP, and the mixture was incubated at 37°C for 3 minutes. 25  $\mu$  of the aggregation inducing substance [adenosine diphosphate (ADP); final concentration 5  $\mu$  or collagen: final concentration 5  $\mu$ g/ml] was added, the mixture was measured for 5 minutes by blood platelet aggregation ability measurement apparatus (Aggricoda TM-PA-3210, Kyoto Dai-ichi Kagaku) to obtain the maximum aggregation rate, and there was calculated the concentration of the test drug (IC $_{\infty}$ ) which brings about 50% inhibition to the maximum aggregation rate obtained by adding the aggregation inducing substance to PRP containing the solvent only.

The compound numbers in Table 1 correspond to those in the Examples described below.

35

10

40

45

50

Table 1

5
•

	Compound	IC <sub>5</sub>	0 (x μM)	Compound	IC <sub>5</sub>	(Mμ x) 0
10	No.	ADP	Collagen	No.	ADP	Collagen
	1	33	14	43	14.0	7.7
	2	28	32	44	10.3	7.3
	- 4	13	16	45	4.4	5.2
15	5	24	23.5	52	7.9	-
	6	24	18	53	4.9	-
	7	12	23	54	. 11.2	15.5
20	8	9.2	13.6	55	10.5	8.3
	9	15	12	56	2.9	6.5
	10	36	26	67	27.7	11.0
25	11	>30	22	68	13.6	7.5
	12	5.6	4.7	75	3.8	5.4
	15	21.5	16.6	76	14.3	10.5
30	16	12.5	4.1	77	4.3	2.9
30	17	7.7	5.0	78	6.2	8.3
	18	6.6	3.2	79	4.3	5.1
	21	30.9		80	7.4	10.9
35	22	41.3	-	81	-5.5	7.0
	24	6.4		85	17.7	14.4
	25	11.1	6.6	86	6.2	5.3
40	26	16.5	9.5	91	9.7	6.7
	29	9.0	8.1	92	7.3	6.5
	32	3.5	3.8	93	18.3	8.7
45	33	11.9	12.5	94	8.0	6.9
_	34	8.2	6.6	95	15.4	2.5
	37	21.2	17.8	96	3.9	3.7
	38	9.0	4.6	97	16.0	3.2
50	39	>30	>30	98	11.2	8.8
	40	11.3	13.2	103	18.5	6.7
	41	4.2	5.1	papaverin	>100	>100
55				_		

Experiment 2 [prolonging test of the bleeding time in mouse]

Six male ICR strain mice weighing 20 g for each group were administered orally with 300 mg/kg of the test drug (all the test drugs were used in the form of the suspension in 0.5% CMC). Two hours after administration, 5 mm of the tail from the top was cut under pentobarbital anesthesia, and the bleeding was observed by tapping at the cutting site with a filter paper every 15 seconds. The time when the bleeding was observed stopping for one minute is defined as the arrest point of bleeding, and the duration required from the time when the cutting was done to the arrest point of bleeding is defined as the bleeding time. The observation was carried out up to 1200 seconds. Ticlopidine was used as a positive control.

The results were shown in Table 2. The compound numbers in Table 2 correspond to those in the Examples described below.

Table 2

15

20

25

Compound No.	Bleeding time ± standard error
53	270.0 ± 54.08
56	277.5 ± 36.90
ticlopidine	1127.5 ± 72.50 (note)
the solvent	305.0 ± 77.23

(Note) P < 0.05 by Mann and Whitney's U test.

30

35

The following Examples illustrate the method for preparing the compound of the present invention in more detail.

### Example 1

## Preparation of (E)-3-(2-oxopropylidene)-1,4-diphenyl-2-azetidinone (Compound 1)

To a solution of 0.67 g of acetylmethylene triphenylphosphorane in 70 ml of benzene was added at room temperature under a nitrogen atmosphere a solution of 0.50 g of 1,4-diphenyl-2,3-azetidinedione in 30 ml of benzene, and the mixture was stirred overnight. After completion of the reaction, the benzene was evaporated, and the residue was applied to silica gel column chromatography (eluent; methylene chloride). The desired fractions were combined, the solvent was evaporated, and the residue was recrystallized from ethanol to give the title compound as pale yellow needles. Yield 0.32 g, m.p. 157.5 - 158.5°C

#### Example 2

Following the similar procedure of that of Example 1, there were obtained the compounds 2 to 118, which were listed in Table 3 including the compound obtained in Example 1.

5			m.p. (°C)	157.5-158.5	149-150.5	130.5-132.5	226-227	174-177	227.5-228.5	147.5-150	222-223	239.5-241	250.5-256
15 20			R2	methyl	ethyl	ethoxy	phenyl	p-methylphenyl	p-methoxyphenyl	o,p-dimethoxy- phenyl	p-fluorophenyl	p-chlorophenyl	p-bromophenyl
25	Table 3	(x) <sub>g</sub>	H	u	¥	Ð	<u> </u>	<b>4</b>	14	0 4	Ç <b>i</b>	<u>p</u>	<u>Di</u>
35 ` 40			$^{\mathrm{R}1}$	phenyl	phenyl	phenyl	phenyl	phenyl	phenyl	phenyl	phenyl	phenyl	phenyl
45 50		·	x(x)	_	_		_						
55			Compound (	1 н	2 н	æ e	4 H	5	н 9	7 н	Н 8	н 6	10 H

55	50	45	: 40	35	30	25	20	15	10	• 5
				Table	Table 3 (Cont'd)	(p.				-
11	E		phenyl				p-biphenyl	nyl	250-250.	ر. ا
12	ш		phenyl				p-nitrophenyl	phenyl	235.5-236.5	9.9
13	ш		phenyl				amino		212-213	
14	H		phenyl		•		l-adamantyl	ntyl	198.5-200	0
15	=		phenyl				ethoxyc methyl	ethoxycarbonyl- methyl	154.5-159.5	59.5
16	Ħ		o-meth	o-methylphenyl			p-metho	p-methoxyphenyl	142-144	
17	ш		o-meth	o-methylphenyl			p-fluorophenyl	ophenyl	140.4-141.9	11.9
18	ш		o-meth	o-methylphenyl			p-nitrophenyl	phenyl	199.5-200.4	00.4
19	Ħ		2,6-di	2,6-dimethylphenyl	enyl	. •	p-fluorophenyl	opheny1	188-189.5	rύ
20	н	•	2,6-di	2,6-dimethylphenyl	enyl		p-nitrophenyl	phenyl	300 or above	pove
21	н		o-meth	o-methyl-p-chlorophenyl	oropheny.	_	p-methylphenyl	1pheny1	142-144	
22	m		o-meth	o-methyl-p-chlorophenyl	oropheny.	_	p-metho	p-methoxyphenyl	147-148.5	ι,
23	н		o-meth	o-methyl-p-chlorophenyl	oropheny	_	p-fluorophenyl	ophenyl	172-174	•
24	<b>=</b>		o-meth	o-methyl-p-chlorophenyl	oropheny	-	p-nitrophenyl	phenyl	195-196	
25	H		2-meth	2-methyl-5-chlorophenyl	oropheny		methyl		149.5-151.5	1.5

55	50	45	40	Table	8 3 (Cont'd)	25	20	15	5	
26	Ħ		2-methy]	2-methyl-5-chlorophenyl	ophenyl		phenyl		145-147	
27	Ħ		2-methy]	2-methyl-5-chlorophenyl	ophenyl		p-fluorophenyl	phenyl	140-142	
28	H		2-methy]	2-methyl-5-chlorophenyl	ophenyl		p-nitrophenyl	nenyl	195.5-197	
29	H		p-fluorophenyl	ophenyl			phenyl		206-208.5	
30			p-fluorophenyl	ophenyl			p-fluorophenyl	pheny1	211-213	
31	Ħ		p-fluorophenyl	ophenyl			p-chlorophenyl	henyl	221.5-224	
32	H		p-fluorophenyl	phenyl			p-nitrophenyl	nenyl	204.5-207	
33	Ħ		o-fluorophenyl	phenyl			p-fluorophenyl	henyl	180.5-183	
34	ш		o-fluorophenyl	pheny1			p-nitrophenyl	nenyl	219.7-221	
35			o-chlorophenyl	phenyl		. •	p-fluorophenyl	henyl	146-147.5	
36	H		o-chlorophenyl	phenyl			p-nitrophenyl	nenyl	189-191	
37	Ħ		3,5-dick	3,5-dichlorophenyl	ул		p-fluorophenyl	henyl	200.2-201.5	
38	ж		3,5-dich	3,5-dichlorophenyl	уl		p-nitrophenyl		206 (decomposition)	n C
39	Ħ		p-bromophenyl	henyl			p-methoxyphenyl	phenyl	208-209	
40	æ		p-bromophenyl	henyl			p-fluorophenyl	heny1	211.5-213	

55	50	45	40	35	30	25	20	15	10	5
				Table	Table 3 (Cont'd)					
41	Œ		p-bromophenyl	henyl			p-nitrophenyl,	enyl,	222-224	
42	ж		o-methoxyphenyl	(yphenyl			p-nitrophenyl	enyl	219-221.2	7
43	Ħ		m-triflu	m-trifluoromethylphenyl	lphenyl		phenyl		174-177	
44	H		m-triflu	m-trifluoromethylphenyl	1phenyl		p-fluorophenyl	nenyl	159.5-161	_
45	H		m-triflu	m-trifluoromethylphenyl	1pheny1		p-nitrophenyl	enyl	181.5-184	₹'
46	н		p-dimeth	p-dimethylaminophenyl	henyl		p-nitrophenyl	anyl	168-170	
47	H	•	p-carbox	p-carboxylphenyl			p-fluorophenyl	nenyl	300 or above	oove
48	н		p-dichloroacetylphenyl	roacetyl	phenyl		p-fluorophenyl	nenyl	180.5-183.5	3.5
49	H		p-dichloroacetylphenyl	roacetyl	phenyl		p-nitrophenyl	ny1	190.5-192.5	2.5
20	æ		benzyl				methyl		76.5-78.5	,0
51	Ħ		benzyl				phenyl		111.5-113.5	3.5
52	æ		benzyl				p-fluorophenyl	ienyl	105-107.5	10
53	Œ		benzyl				p-nitrophenyl	nyl	122-126	
54	H		o-chlorobenzyl	benzyl		•	methyl		78-79	
55	Ħ		o-chlorobenzyl	benzyl		ı	p-fluorophenyl	enyl	74-76	

55	50	45	40	. 35	30	25	20	15	10	5
				Table 3	(Cont'd)					
26	н		o-chlorobenzyl	)enzyl		Д	p-nitrophenyl	ıny1	113-115	
57	Ħ		l(S)-phenethyl	nethyl		<b>—</b>	p-nitrophenyl	nyl	127.5-130.5	.5
28	Ħ		1-carboxs	1-carboxy-2-phenethyl	thyl	_	p-fluorophenyl	ıenyl	250-255	
59	Ħ		propyl			_	p-fluorophenyl	nenyl	88.5-91	
09	æ		propyl			~	p-nitrophenyl	inyl	127.5-130.5	.5
61	Ħ		cyclohexyl	71		_	methyl		124-127	
62	æ		cyclohexyl	71		_	p-fluorophenyl	lenyl	125-126.5	40
63	н		cyclohexyl	y1			p-nitrophenyl	any 1	199-202.5	10
64	н		l,2-bis(m ethyl	l,2-bis(methoxycarbonyl)- ethyl	rbonyl) –	-	p-fluorophenyl	ıenyl	126-128	
65	p-methyl		phenyl				p-fluorophenyl	enyl	208.5-211	
99	p-methyl		phenyl			_	p-nitrophenyl	any1	240.5-242.	5.5
29	p-ethyl		o-methylphenyl	jhenyl		_	p-fluorophenyl	neny1	143-144.2	•
89	p-ethyl		o-methylphenyl	jheny1		_	p-nitrophenyl	iny1	157.2-158.	3.6
69	o-methoxy		o-methylphenyl	jhenyl		_	p-fluorophenyl	nenyl	133-135.5	
70	o-methoxy		o-methylphenyl	jheny1		_	p-nitrophenyl	ınyı	178-180.5	

	50	45	40	35	30	25	<b>2</b> 0 -	15	10	5
				Table 3	3 (Cont'd)	æ				
	m-methoxy		phenyl				p-fluorophenyl	henyl	173.5-176.2	76.2
	m-methoxy		phenyl				p-nitrophenyl	enyl	194.5-196.5	96.5
	3,4-dimethoxy	hoxy	phenyl				p-fluorophenyl	henyl	164.5-169	69
	3,4-dimethoxy	hoxy	phenyl				p-nitrophenyl	enyl	192-195	
	p-hydroxy		phenyl				p-nitrophenyl	enyl	166.5-167.	67.5
	p-fluoro		phenyl				p-fluorophenyl	heny1	209.5-211	11
	p-fluoro	,	phenyl				p-nitrophenyl	enyl	225-226	
	p-fluoro		o-methylphenyl	phenyl			p-fluorophenyl	henyl	157-159.5	.5
	p-fluoro		o-methylphenyl	phenyl			p-nitrophenyl	enyl	193-195.5	ις
	o-fluoro		phenyl				p-fluorophenyl	henyl	191.3-192.2	92.2
	o-fluoro		phenyl				p-nitrophenyl	ıyı	224.8-226.7	6.7
	o-chloro		phenyl				p-fluorophenyl	nenyl	213.5-216	9 1
	p-chloro		o-methylphenyl	phenyl			p-fluorophenyl	nenyl	150-151.5	Ŋ
	p-chloro		o-methylphenyl	pheny1		-	p-nitrophenyl	nv.]	180-182	
,	p-bromo		o-methylphenyl	phenyl			p-fluorophenyl	enyl	157.4-158.7	18.7

55	50	45	40	35	30	25	20	75	10	5
				Table	Table 3 (Cont'd)	<b>a</b>				
98	p-bromo		o-methy	o-methylphenyl			p-nitrophenyl	heny1	180-180.5	0.5
87	o-bromo		phenyl				p-fluorophenyl	phenyl	225-227	7
88	o-promo		phenyl				p-nitrophenyl	henyl	210-212	2
89	p-cyano		o-methy	o-methylphenyl			p-fluorophenyl	phenyl	182.2-187.7	187.7
06	p-cyano		o-methy	o-methylphenyl			p-nitrophenyl	henyl	180.5-183.7	183.7
91	Ħ		p-methy	p-methylbenzyl			p-nitrophenyl	henyl	147-148	œ
92	Ħ		p-metho	p-methoxylbenzyl	-		p-nitrophenyl	henyl	110-112	2
93	Ħ		p-fluor	p-fluorobenzyl			p-nitrophenyl	henyl	156.5-158.	158.5
94	н		o-metho	o-methoxybenzyl			p-nitrophenyl	henyl	146.5-148.5	148.5
95	н		o-trif1	o-trifluoromethylbenzyl	ylbenzyl		p-nitrophenyl	henyl	126-127.5	7.5
96	Ħ		o-fluor	o-fluorobenzyl			p-nitrophenyl	henyl	116-117	7
16	Œ		m-chlor	m-chlorobenzyl			p-nitrophenyl	henyl	145-147	7
86	Ħ		p-chlor	p-chlorobenzyl			p-nitrophenyl	henyl	157.5-159.5	159.5
66	Ħ		m-trifl	<pre>m-trifluoromethylbenzyl</pre>	ylbenzyl		p-nitrophenyl	henyl	124-126	9
100	E		p-trifl	p-trifluoromethylbenzyl	ylbenzyl		p-nitrophenyl	henyl	107.5-109	109

55	<b>4</b> 5	40	35	30	25	20	15	10	_
			Table	Table 3 (Cont'd)	Œ.				
101		m-methoxybenzyl	ybenzyl			p-nitrophenyl	henyl	124-126	
102	н	3,4-meth	ylenedio	3,4-methylenedioxybenzyl		p-nitrophenyl	henyl	148-151	
103	н	2,4-dichlorobenzyl	lorobenz	уl		p-nitrophenyl	henyl	86-96	
104	н	3,4-dichlorobenzyl	lorobenz	:41		p-nitrophenyl	henyl	145.5-148	_
105	н	l-naphthylmethyl	ylmethyl			p-nitrophenyl	henyl	167.5-169	_
901	H	o-fluorobenzyl	benzyl			p-fluorophenyl	phenyl	96-97.5	
107	н	m-methoxybenzyl	ybenzyl			p-fluorophenyl	phenyl	108-110.5	
108	Ħ	<pre>m-trifluoromethylbenzyl</pre>	oromethy	lbenzyl		p-fluorophenyl	phenyl	100-102	
109	Ħ	p-trifluoromethylbenzyl	oromethy	lbenzyl		p-fluorophenyl	phenyl	136-138	
110	Ħ	3,4-dichlorobenzyl	lorobenz	yı		p-fluorophenyl	phenyl	111-113	
111	o-methyl	benzyl				p-nitrophenyl	henyl	111-114	
112	p-methoxy	benzyl				p-nitrophenyl	henyl	127-128	
113	p-fluoro	benzyl				p-nitrophenyl	henyl	118-120	
114	m-chloro	benzyl				p-nitrophenyl	henyl	82-87	
115	p-fluoro	o-chlorobenzyl	benzyl			p-nitrophenyl	henyl	98.5-101.5	S

10	·	155-156	153.5-157	115.5-121.	
15		p-nitrophenyl	p-nitrophenýl	p-nitrophenyl	
20		u-d	u-d	u-đ	
<b>25</b>	ont'd)			,	
30	Table 3 (Cont'd)	ул	уl	۲٦	
<b>35</b>	Tab	o-chlorobenzyl	o-chlorobenzyl	o-chlorobenzyl	
		ey l	•	,	!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!
45		p-isoprop	o-fluoro	p-trifluoro- methyl	1
55		116	117	118	

Claims

5

10

15

25

35

1. 2-Azetidinone derivatives represented by the following formula

$$\begin{array}{c}
\mathbb{R}^{2} \\
\mathbb{R}^{2}
\end{array}$$

wherein X is a hydrogen atom, a halogen atom, a lower alkyl group, a lower alkoxy group, a hydroxyl group or a cyano group, 1 is 1 or 2, R1 is a lower alkyl group, a cycloalkyl group, a 1-naphthylmethyl group, a 1-phenethyl group, 1-carboxy-2-phenethyl group, a group of the formula

(wherein Y and Y' are the same or different and each is a hydrogen atom, a halogen atom, a lower alkyl group, a lower alkoxy group, a dimethylamino group, a carboxyl group, a dichloroacetyl group or a trifluoromethyl group, or Y and Y' together form a methylenedioxy group, and m is 0 or 1) or a group of the formula

(wherein R<sup>2</sup> is a lower alkyl group), and R<sup>2</sup> is a lower alkyl group, a lower alkoxy group, an amino group, an adamantyl group, a lower alkoxycarbonylmethyl group, or a group of the formula

(wherein Z is a hydrogen atom, a halogen atom, a lower alkyl group, a lower alkoxy group, a phenyl group or a nitro group, and n is 1 or 2).

2. Blood platelet aggregation inhibiting agents containing 2-azetidinone derivatives represented by the general formula

55

$$R^2$$
  $O$   $(X)_{\ell}$ 

10

wherein X is a hydrogen atom, a halogen atom, a lower alkyl group, a lower alkoxy group, a hydroxyl group or a cyano group, t is 1 or 2, R<sup>1</sup> is a lower alkyl group, a cycloalkyl group, a 1-naphthylmethyl group, a 1-phenethyl group, 1-carboxy-2-phenethyl group, a group of the formula

15

20

(wherein Y and Y' are the same or different and each is a hydrogen atom, a halogen atom, a lower alkyl group, a lower alkoxy group, a dimethylamino group, a carboxyl group, a dichloroacetyl group or a trifluoromethyl group, or Y and Y' together form a methylenedioxy group, and m is 0 or 1) or a group of the formula

30

25

35

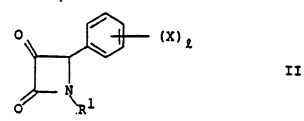
(wherein R<sup>2</sup> is a lower alkyl group), and R<sup>2</sup> is a lower alkyl group, a lower alkoxy group, an amino group, an adamantyl group, a lower alkoxycarbonylmethyl group, or a group of the formula

40

(wherein Z is a hydrogen atom, a halogen atom, a lower alkyl group, a lower alkoxy group, a phenyl group or a nitro group, and n is 1 or 2).

- 3. A 2-azetidinone derivative according to Claim 1, wherein the oxyalkylidene substituent has the E-configuation.
- 4. A 2-azetidinone derivative according to Claim 1 or Claim 3, wherein the configuration due to the asymmetric carbon atom at the 4-position is of the dl-form.
- 5. A process for producing a 2-azetidinone derivative of the formula given and defined in Claim 1, which comprises reacting a compound of the formula

50



wherein R1, X and 1 are as defined in Claim 1, with a Wittig reagent of the formula

$$\mathbb{R}^2$$
  $\longrightarrow \mathbb{P}(\mathbb{C}_6^{\mathbb{H}_5})_3$ 

wherein R2 is as defined in Claim 1.

- 6. A 2-azetidinone derivative of the formula given and defined in Claim 1 for use as a pharmaceutical.
- 7. A 2-azetidinone derivative of the formula given and defined in Claim 1 for use as a blood platelet aggregation inhibiting agent.
- 8. A pharmaceutical composition comprising a 2-azetidinone derivative of the formula given and defined in Claim 1 and a pharmaceutically acceptable diluent or carrier.



# **EUROPEAN SEARCH REPORT**

	DOCUMENTS CONSI	EP 87308942.9			
ategory		indication, where appropria int passages	ie,	Relevant to claim	CLASSIFICATION OF THE APPLICATION (Int. Cl.4)
Α	TETRAHEDRON, vol	. 41, no. 2,	1985 1	,3-5	C 07 D 205/08
	MORI et al.: "Ne β-lactames" pages 375-385	•	ì	,	A 61 K 31/395
		385 (compound 20c, 20c') *	ls.		
A	LIEBIGS ANNALEN DER CHEMIE, 1983, Heft 5			, 3–5	
	HH. OTTO et al und Stereochemie benzyl)-1,4-diph pages 1152-1168	e von 3-( <b>≪-</b> Hyc	lroxy-		
	* Pages 1153; 3,5); pages pounds 4,41	, 1158 (compou s 1165-1168 (c f,8) *	eom-		
A	ARCHIV DER PHARMAZIE, vol. 319,		.9. 1	1,3-5 TECHNICAL FIELDS SEARCHED (int. Cl.4)	
	no. 3, March 1986			,,,,,	C 07 D 205/00
	BERGMANN et al.: "Zur N- und C- Silylierung von β-Lactamen" pages 203-216				A 61 K 31/00
	* Pages 208,2 14,15) *	214,215 (compo	ounds		
A	EP - A1 - 0 149 PHARM.)	419 (NIPPON 2	OKI 2	2,6-8	
i	* Page 1, las 2; claims 1	st two lines; L5-19 *	page		
	•				:
<del></del> _	The present search report has b	een drawn up for all claims			
Place of search Date of comple		Date of completion of	the search		Examiner
	VIENNA	07-01-1988	3		JANISCH
Y : pa	CATEGORY OF CITED DOCL inticularly relevant if taken alone tricularly relevant if combined w ocument of the same category chnological background in-written disclosure	E :	theory or prin earlier patent after the filing document cit document cit	document date ed in the a	orlying the invention t, but published on, or pplication or reasons

# **EUROPEAN SEARCH REPORT**

. Application number

Eategory	Citation of document	with indication, where appropriate,	0.1	
n 4		levent passages	Relevant to claim	
D,A	TETRAHEDRON L	ETTERS, vol. 25, n	.0. 5	
	MANHAS et al. synthesis of a pages 4733-6	: "A convenient azetidine-2,3-dion	es"	
	* Page 4735	5 *		
				·
		·		i
ł				
				TECHNICAL FIELDS
				SEARCHED (Int. Cl.4)
-		•		
	The expect ease to		_	
The present search report has been drawn up for all claims  Place of search  Date of completion of the search				
VIENNA		Date of completion of the search 07-01-1988	'	Examiner
07-01-1388				JANISCH invention
			itent document	. but published on, or

EPO form 1503 03 62

